

Figure 1

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ion trans: domain 1 of 1, from 472 to 661: score 0.1, E = 1.2
    (SEQ ID NO:9) *->ilfildllfvllflleivlkfiayglkstsniaakylksifnildll
                     ++ i + ++f++ ++1
                                        ++++
                                                   ++ y +++
                     DWRIAMTYERIFFICLEILVCAIHPI-----PGNYTFTWTA-RLAF 511
    52906
              472
                  ailplllllvlflsgteqvakkrlrerfslelsqwyyrilrflrlLrllR
              512 SYAPS--TTTADVDII------LSIPMFLRLYL 536
    52906
                  lLrllrllrrletlf.e....fe.lgtlaWslqslgralksilrfllll
                     + r++ ++ +lf+ ++++++ 1 ++
              537 --- IARVMLLHSKLFtDtssrsIGaLNKI---NFNTRFVMKTLMTICPGT 580
    52906
                  lllligfsvigyllfkgyedlsenevdgnsefssyfdafyflfvtlttvG
                  +ll++ s+ ++++ + + +e+ d+ + s++ a++ +t++++G
    52906
              581 VLLVF--SISLWIIAAWTVRACERYHDQQDVTSNFLGAMWLISITFLSIG 628
                  fGdlvpvwlgiiffvlffiivgllllnlliavi<-*
                  +Gd+vp++++ ++l+ i+g ++l +av+
    52906
              629 YGDMVPNTYCGKGVCLLTGIMGAGCTALVVAVV
                                                     661
```

Figure 2

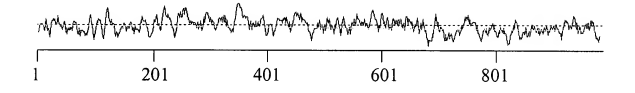


Figure 3

	<pre>1 of 1, from 247 to 467: score 94.8, E = 1.7e-24 *->ilfildllfvllflleivlkfiayglkstsniaak +l ld + +++fl++ivl+f+ + + ++++ ++++ i++</pre>	
33408 247	WLV-LDSVVDVIFLVDIVLNFH-TTFVgpggevisdpklIRMN	287
33408 288	ylksifnildlailplllllvlflsgteqvakkrlrerfslelsqwyyr ylk++f +dll++lp+++ ++ ++ ++ YLKTWFV-IDLLSCLPYDIINAFENVDEG	315
	<pre>ilrflrlLrllRlLrllrllrrletlfefelgtlaWslqslg.ralksil i +++++L+ +R</pre>	0.10
33408 316	ISSLFSSLKVVRLLRLGRVARKLDHYLEYGAaVLVLLVC	354
	rflllllllligfsvigyllfkgyedlse f+l++++l ++++ ig + + ++ + +++ +++++++++++	
33408 355	VFGLVAHWLACIWYSIGDYEVIDEVTNTIqidswlyqlalsigtpyrynt	404
33408 405	nevdgnsefssyfdafyflfvtlttvGfGdlvpv.wlgiiffvlffi + + +++g+s+ s y ++yf++++ltt+GfG++ p++ +++f v++++ sagIWEGGPSKDSLYVSSLYFTMTSLTTIGFGNIAPTtDVEKMFSVAMMM	454
33400 403		434
33408 455	<pre>ivgllllnlliavi<-* +++ ll ++++++ VGS-LLYATIFGNV 467</pre>	

Figure 4A

Figure 4B

33408 MPGGKRGLVA PQNTFLENIV RRSSESSFLL GNAQIVDWPV VYSNDGFCKL rat EAG2 MPGGKRGLVA PONTFLENIV RRSSESSFLL GNAOIVDWPV VYSNDGFCKL (SEQ ID NO:12) SGYHRADVMQ KSSTCSFMYG ELTDKKTIEK VRQTFDNYES NCFEVLLYKK SGYHRADVMQ KSSTCSFMYG ELTDKKTIEK VRQTFDNYES NCFEVLLYKK NRTPVWFYMQ IAPIRNEHEK VVLFLCTFKD ITLFKOPIED DSTKGWTKFA NRTPVWFYMQ IAPIRNEHEK VVLFLCTFKD ITLFKOPIED DSTKGWTKFA RLTRALTNSR SVLQQLTPMN KTEVVHKHSR LAEVLQLGSD ILPQYKQEAP RLTRALTNSR SVLQQLTPMN KTETVHKHSR LAEVLQLGSD ILPQYKQEAP KTPPHIILHY CAFKTTWDWV ILILTFYTAI MVPYNVSFKT KQNNIAWLVL KTPPHIILHY CAFKTTWDWV ILILTFYTAI MVPYNVSFKT KQNNIAWLVL DSVVDVIFLV DIVLNFHTTF VGPGGEVISD PKLIRMNYLK TWFVIDLLSC DSVVDVIFLV DIVLNFHTTF VGPGGEVISD PKLIRMNYLK TWFVIDLLSC LPYDIINAFE NVDEGISSLF SSLKVVRLLR LGRVARKLDH YLEYGAAVLV LPYDIINAFE NVDEGISSLF SSLKVVRLLR LGRVARKLDH YLEYGAAVLV LLVCVFGLVA HWLACIWYSI GDYEVIDEVT NTIQIDSWLY QLALSIGTPY LLVCVFGLVA HWLACIWYSI GDYEVIDEVT NTIQIDSWLY OLALSIRTPY RYNTSAGIWE GGPSKDSLYV SSLYFTMTSL TTIGFGNIAP TTDVEKMFSV RYNTSAGIWE GGPSKDSLYV SSLYFTMTSL TTIGFGNIAP TTDVEKMFSV AMMMVGSLLY ATIFGNVTTI FQQMYANTNR YHEMLNNVRD FLKLYQVPKG AMMMVGSLLY ATIFGNVTTI FQQMYANTNR YHEMLNNVRD FLKLYQVPKG LSERVMDYIV STWSMSKGID TEKVLSICPK DMRADICVHL NRKVFNEHPA LSERVMDYIV STWSMSKGID TEKVLSICPK DMRADICVHL NRKVFNEHPA FRLASDGCLR ALAVEFQTIH CAPGDLIYHA GESVDALCFV VSGSLEVIQD FRLASDGCLR ALAVEFQTIH CAPGDLIYHA GESVDALCFV VSGSLEVIQD DEVVAILGKG DVFGDIFWKE TTLAHACANV RALTYCDLHI IKREALLKVL EEVVAILGKG DVFGDIFWKE TTLAHACANV RALTYCDLHI IKREALLKVL

Figure 4C

DFYTAFANSF SRNLTLTCNL RKRIIFRKIS DVKKEEERL RQKNEVTLSI
DFYTAFANSF SRNLTLTCNL RKRIIFRKIS DVKKEEERL RQKNEVTLSI
PVDHPVRKLF QKFKQQKELR NQGSTQGDPE RNQLQVESRS LQNGTSITGT
PVDHPVRKLF QKFKQQKELR NQGSAQSDPE RSQLQVESRP LQNGASITGT

SVVTVSQITP IQTSLAYVKT SESLKQNNRD AMELKPNGGA DQKCLKVNSP
SVVTVSQITP IQTSLAYVKT SETLKQNNRD AMELKPNGGA EPKCLKVNSP

IRMKNGNGKG WLRLKNNMGA HEEKKEDWNN VTKAESMGLL SEDPKSSDSE
IRMKNGNGKG WLRLKNNMGA HEEKKEEWNN VTKAESMGLL SEDPKGSDSE

NSVTKNPLRK TDSCDSGITK SDLRLDKAGE ARSPLEHSPI QADAKHPFYP
NSVTKNPLRK TDSCDSGITK SDLRLDKAGE ARSPLEHSPS QADAKHPFYP

IPEQALQTTL QEVKHELKED IQLLSCRMTA LEKQVAEILK ILSEKSVPQA
IPEQALQTTL QEVKHELKED IQLLSCRMTA LEKQVAEILK LLSEKSVPQT

SSPKSQMPLQ VPPQIPCQDI FSVSRPESPE SDKDEIHF
SSPKPQIPLQ VPPQIPCQDI FSVSRPESPE SDKDEIHF

Figure 4D

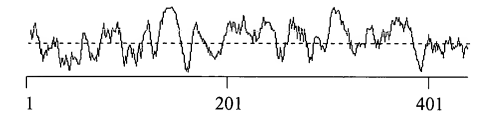


Figure 5

		f 1, from 3 to 101: score 169.0, E = 7.9e-47
(SEQ ID NO	0:11)	*->ErvrLNVGGkrFeTsksTLtrfkpdTlLgrllktdsdvhearlrlcd
		Er++LNV G+rFeT+++TL rf pdTlLg++++ r ++
12189	3	ERLVLNVAGLRFETRARTLGRF-PDTLLGDPARR-GR 37
		fyddetgEyFFDRsPkhFetILnfYRtGdGkLhrp.evcldsfleEleFy
		fydd++ EyFFDR++++F+++L++Y++G G+L+rp +v+ld+fleE +Fy
12189	38	FYDDARREYFFDRHRPSFDAVLYYYQSG-GRLRRPaHVPLDVFLEEVAFY 86
		gldelaiesCcedeY<-*
		gl+ a++ +ede+
12189	87	GLGAAALARLREDEG 101

Figure 6A

```
(SEQ ID NO:9) *->ilfildllfvllflleivlkfiayglkstsniaakylksifnildll
                ++f++++l++ +f +e+++++ ++ k
                                                 a ++k+++n++d+
12189
         198
                PFFVVETLCICWFSFELLVRLLVCPSK-----AIFFKNVMNLIDFV 238
             ailplllllvlflsgteqvakkrlrerfslelsqwyyrilrflrlLrllR
             ailp+++ l+ l+++
                                                 ++++ + +L +lR
          239 AILPYFVALGTELARQ-------RGVGQQAMSLAILR 268
12189
              lLrllrllrrletlfefelgtlaWslqslg.ralksilrfllllllllig
              ++rl+r++r ++ +
                               +++
                                      lq+lg+++ +s+ ++ll+++l+ig
          269 VIRLVRVFRIFKLSR---HSKG---LQILGqTLRASMRELGLLIFFLFIG 312
12189
             fsvigyllfkgyedlsenevdgnsefssyfdafyflfvtlttvGfGdlvp
                                   +s f+s++++f+++vt+ttvG+Gd+ p
              + +++ ++++ + d+
         313 VVLFSSAVYFAEVDRV-----DSHFTSIPESFWWAVVTMTTVGYGDMAP 356
12189
             v.wlgiiffvlffiivgllllnlliavi<-*
             v+++g+i++ ++++i+g+l+++l+++vi
12189
         357 VtVGGKIVG-SLCAIAGVLTISLPVPVI
                                             383
```

ion trans: domain 1 of 1, from 198 to 383: score 144.8, E = 1.5e-39

Figure 6B

Mouse Kv1.7

12189

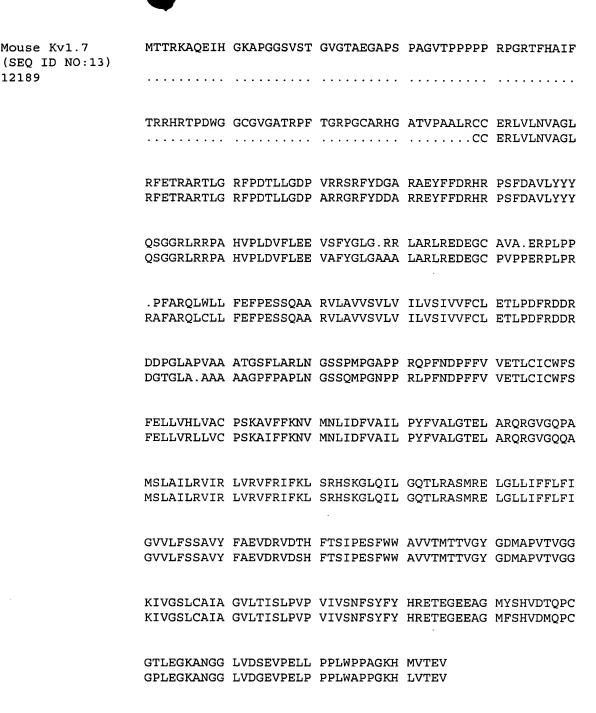


Figure 6C